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# OM protein - protein search, using sw model

Run on: June 25, 2003, 14:20:41 ; Search time 33 Seconds  
(without alignments)  
444.169 Million cell updates/sec

Title: US-09-622-613b-24

Perfect score: 601  
Sequence: 1 SWMAPFOCKHIINTPICNT.....ICVCKENQYVHFAGIGRCP 110

Scoring table: BL/SUM62  
Gapop 10.0 , Gapext 0.5

Searched: 903470 seqs, 133250620 residues

Total number of hits satisfying chosen parameters: 908470

Minimum DB seq length: 0  
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%  
Maximum Match 100%

Listing first 45 summaries

## Database :

A.Geneseq\_101002:\*

- 1: /SID52/gcgdata/geneseq/geneseq-emb1/AA1980.DAT:\*
- 2: /SID52/gcgdata/geneseq/geneseq-emb1/AA1981.DAT:\*
- 3: /SID52/gcgdata/geneseq/geneseq-emb1/AA1982.DAT:\*
- 4: /SID52/gcgdata/geneseq/geneseq-emb1/AA1983.DAT:\*
- 5: /SID52/gcgdata/geneseq/geneseq-emb1/AA1984.DAT:\*
- 6: /SID52/gcgdata/geneseq/geneseq-emb1/AA1985.DAT:\*
- 7: /SID52/gcgdata/geneseq/geneseq-emb1/AA1986.DAT:\*
- 8: /SID52/gcgdata/geneseq/geneseq-emb1/AA1987.DAT:\*
- 9: /SID52/gcgdata/geneseq/geneseq-emb1/AA1988.DAT:\*
- 10: /SID52/gcgdata/geneseq/geneseq-emb1/AA1989.DAT:\*
- 11: /SID52/gcgdata/geneseq/geneseq-emb1/AA1990.DAT:\*
- 12: /SID52/gcgdata/geneseq/geneseq-emb1/AA1991.DAT:\*
- 13: /SID52/gcgdata/geneseq/geneseq-emb1/AA1992.DAT:\*
- 14: /SID52/gcgdata/geneseq/geneseq-emb1/AA1993.DAT:\*
- 15: /SID52/gcgdata/geneseq/geneseq-emb1/AA1994.DAT:\*
- 16: /SID52/gcgdata/geneseq/geneseq-emb1/AA1995.DAT:\*
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- 18: /SID52/gcgdata/geneseq/geneseq-emb1/AA1997.DAT:\*
- 19: /SID52/gcgdata/geneseq/geneseq-emb1/AA1998.DAT:\*
- 20: /SID52/gcgdata/geneseq/geneseq-emb1/AA1999.DAT:\*
- 21: /SID52/gcgdata/geneseq/geneseq-emb1/AA2000.DAT:\*
- 22: /SID52/gcgdata/geneseq/geneseq-emb1/AA2001.DAT:\*
- 23: /SID52/gcgdata/geneseq/geneseq-emb1/AA2002.DAT:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	601	100.0	110	20	AA128877
2	601	100.0	111	20	AA128878
3	597	99.3	110	20	AA128872
4	597	99.3	111	20	AA128873
5	591	99.3	110	20	AA128874
6	591	99.3	111	20	AA128876
7	586.5	97.6	111	20	AA128877
8	282.5	47.0	104	18	AA128878
9	280.5	46.7	104	20	AA128879
10	280.5	46.7	105	20	AA128871

11	276.5	46.0	104	20	AA128865	Rana pipiens liver
12	276.5	46.0	105	20	AA128867	Recombinant Met(-1
13	276.5	46.0	112	18	AA128868	R. pipiens recombi
14	276.5	46.0	127	20	AA128879	Rana pipiens Clone
15	276.5	46.0	251	18	AA128880	R. pipiens recombi
16	276.5	46.0	254	18	AA128881	R. pipiens recombi
17	276.5	46.0	355	18	AA128882	R. pipiens recombi
18	276.5	46.0	355	18	AA128883	R. pipiens recombi
19	276.5	46.0	355	18	AA128884	R. pipiens recombi
20	276.5	46.0	355	18	AA128885	R. pipiens recombi
21	273.5	45.5	105	20	AA128886	Recombinant RAPRI
22	272.5	45.3	104	12	AA128887	Recombinant Met(-1
23	272.5	45.3	104	12	AA128888	Protein with activ
24	272.5	45.3	104	17	AA128889	ONCONASE (pharmace
25	272.5	45.3	104	17	AA128890	Protein derived fr
26	272.5	45.3	104	18	AA128891	Recombinant onc pr
27	272.5	45.3	104	18	AA128892	Antitumor protein
28	272.5	45.3	104	18	AA128893	Onconase (RTM) pro
29	272.5	45.3	104	20	AA128894	Frog onconase prot
30	272.5	45.3	104	20	AA128895	Rana pipiens RNase
31	272.5	45.3	104	22	AA128896	Amino acid sequenc
32	272.5	45.3	105	18	AA128897	R. pipiens recombi
33	272.5	45.3	106	18	AA128898	Recombinant frog O
34	272.5	45.3	107	18	AA128899	R. pipiens recombi
35	272.5	45.3	107	18	AA128900	R. pipiens recombi
36	272.5	45.3	107	18	AA128901	R. pipiens recombi
37	272.5	45.3	107	18	AA128902	R. pipiens recombi
38	271.5	45.2	105	18	AA128903	R. pipiens recombi
39	269.5	44.8	104	18	AA128904	Recombinant onc pr
40	267.5	44.5	104	22	AA128905	Amino acid sequenc
41	267.5	44.5	105	18	AA128906	R. pipiens recombi
42	267.5	44.5	105	18	AA128907	R. pipiens recombi
43	267.5	44.5	105	18	AA128908	R. pipiens recombi
44	265.5	44.2	104	18	AA128909	Antitumor generic
45	253.5	42.2	107	18	AA128910	R. pipiens recombi

## ALIGNMENTS

RESULT 1	
AA128877	
ID	AA128877 standard; Protein: 110 AA.
XX	
AC	AA128877;
XX	
DT	25-JAN-2000 (first entry)
XX	
DE	Recombinant RacOR1 Gln1Ser amino acid sequence.
XX	
KW	Recombinant Rana catesbeiana oocyte ribonuclease; RacOR1 Gln1Ser; CD22;
KW	covalently bound; IL2 antibody; ligand binding moiety; cancerous B cell;
KW	bulfrog; Kaposi's sarcoma; human chorionic gonadotropin; hCG; RNase;
KW	signal peptide; recombinant ribonuclease; cytotoxic fusion protein;
KW	cancer; autoimmune disease.
XX	
OS	Rana catesbeiana.
XX	
XX	Synthetic.
FT	
FT	Key
FT	Misc-difference 1
FT	Location/Qualifiers
XX	/note="Wild type Gln replaced with Ser"
PN	W09950398-A2.
XX	
PD	07-OCT-1999.
XX	
PF	26-MAR-1999; 99WO-US06641.
XX	
PR	27-MAR-1998; 98US-0079751.
XX	
PA	(USSH ) US DEPT HEALTH & HUMAN SERVICES.
XX	

PI Newton DL, Rybak SM;  
XX  
XX MPI: 1999-610847/52.  
DR N-PSDB; AAZ08134.  
XX  
PT New recombinant ribonucleases, used for killing target cells, e.g. for  
PT treating cancers, viral infections or autoimmune diseases  
XX  
PS Claim 22; Page 67; 71pp; English.  
XX  
CC The present sequence is a recombinant Rana catesbeiana oocyte  
CC ribonuclease (RACOR1) protein with Gln1Ser. Carboxy terminal end of  
CC recombinant RACOR1 has a covalently bound ligand binding moiety, which  
CC can be a LL2 antibody directed against CD22 on cancerous B cells or  
CC human chorionic gonadotropin (hCG) effective against Kaposi's sarcoma  
CC cells. Recombinant ribonucleases can be expressed in bacteria without an  
CC N-terminal methionine due to the presence of a signal peptide that is  
CC cleaved by bacteria. The soluble expression of ribonuclease allows the  
CC proteins to be fused in-frame with ligand binding moieties to form  
CC cytotoxic fusion proteins. They can be used for treatment of cancer and  
CC autoimmune diseases.  
XX  
SQ Sequence 110 AA;  
Query Match 100.0%; Score 601; DB 20; Length 110;  
Best Local Similarity 100.0%; Pred. No. 3.1e-61;  
Matches 110; Conservative 0; Mismatches 0; Indels 0; Caps 0;  
QY 1 SNNATFOQKHIIPTPIICNTIMDNNIYVGGCKRVNTFISSATVTKATCTGVINMNL 60  
DB 1 SNNATFOQKHIIPTPIICNTIMDNNIYVGGCKRVNTFISSATVTKATCTGVINMNL 60  
QY 61 STTRFQNLNCTRTSITPRCPYSSRTETNYICVCKENQYVPHFAGIGRCP 110  
DB 61 STTRFQNLNCTRTSITPRCPYSSRTETNYICVCKENQYVPHFAGIGRCP 110  
RESULT 2  
AAZ28878  
ID AAY28878 standard; Protein; 111 AA.  
XX  
AC AAY28878;  
XX  
DT 25-JAN-2000 (first entry)  
XX  
DE Recombinant Met(-1) RACOR1 Gln1Ser amino acid sequence.  
XX  
KW Recombinant Met(-1) Rana catesbeiana oocyte ribonuclease Gln1Ser; RACOR1;  
KW covalently bound; LL2 antibody; ligand binding moiety; cancerous B cell;  
KW Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide;  
KW recombinant ribonuclease; cytotoxic fusion protein; cancer; bullfrog;  
KW CD22; RNase; autoimmune disease.  
XX  
OS Rana catesbeiana.  
OS Synthetic.  
OS  
FH Key Location/Qualifiers  
FT MISC-difference 1 /note= "Met not found in wild type RACOR1"  
FT MISC-difference 2 /note= "Wild type Gln replaced with Ser"  
FT  
XX MO9950398-A2.  
XX  
XX 07-OCT-1999.  
XX  
XX 26-MAR-1999; 99WO-US06641.  
XX  
XX 27-MAR-1998; 98US-0079751.  
XX  
XX (USSH ) US DEPT HEALTH & HUMAN SERVICES.  
XX  
XX Newton DL, Rybak SM;  
XX

XX  
XX MPI: 1999-610847/52.  
DR N-PSDB; AAZ08135.  
XX  
PT New recombinant ribonucleases, used for killing target cells, e.g. for  
PT treating cancers, viral infections or autoimmune diseases  
XX  
PS Claim 22; Page 68; 71pp; English.  
XX  
CC The present sequence is a recombinant Rana catesbeiana ribonuclease  
CC (RACOR1) protein with Met at position 1 and Gln2Ser. Carboxy terminal end  
CC of recombinant RACOR1 has a covalently bound ligand binding moiety, which  
CC can be a LL2 antibody directed against CD22 on cancerous B cells or human  
CC chorionic gonadotropin (hCG) effective against Kaposi's sarcoma cells.  
CC Recombinant ribonucleases can be expressed in bacteria without an N-  
CC terminal methionine due to the presence of a signal peptide that is  
CC cleaved by bacteria. The soluble expression of ribonuclease allows the  
CC proteins to be fused in-frame with ligand binding moieties to form  
CC cytotoxic fusion proteins. They can be used for treatment of cancer and  
CC autoimmune diseases.  
XX  
SQ Sequence 111 AA;  
Query Match 100.0%; Score 601; DB 20; Length 111;  
Best Local Similarity 100.0%; Pred. No. 3.1e-61;  
Matches 110; Conservative 0; Mismatches 0; Indels 0; Caps 0;  
QY 1 SNNATFOQKHIIPTPIICNTIMDNNIYVGGCKRVNTFISSATVTKATCTGVINMNL 60  
DB 2 SNNATFOQKHIIPTPIICNTIMDNNIYVGGCKRVNTFISSATVTKATCTGVINMNL 61  
QY 61 STTRFQNLNCTRTSITPRCPYSSRTETNYICVCKENQYVPHFAGIGRCP 110  
DB 62 STTRFQNLNCTRTSITPRCPYSSRTETNYICVCKENQYVPHFAGIGRCP 111  
RESULT 3  
AAZ28872  
ID AAY28872 standard; Protein; 110 AA.  
XX  
AC AAY28872;  
XX  
DT 25-JAN-2000 (first entry)  
XX  
DE Rana catesbeiana oocyte ribonuclease (RACOR1) amino acid sequence.  
XX  
KW Rana catesbeiana oocyte ribonuclease; RACOR1; covalently bound; CD22;  
KW LL2 antibody; ligand binding moiety; cancerous B cell; Kaposi's Sarcoma;  
KW human chorionic gonadotropin; hCG; recombinant ribonuclease; bullfrog;  
KW signal peptide; cytotoxic fusion protein; cancer; autoimmune disease;  
KW RNase.  
XX  
OS Rana catesbeiana.  
OS Synthetic.  
OS  
FH Key Location/Qualifiers  
FT MISC-difference 1 /note= "Met not found in wild type RACOR1"  
FT MISC-difference 2 /note= "Wild type Gln replaced with Ser"  
FT  
XX MO9950398-A2.  
XX  
XX 07-OCT-1999.  
XX  
XX 26-MAR-1999; 99WO-US06641.  
XX  
XX 27-MAR-1998; 98US-0079751.  
XX  
XX (USSH ) US DEPT HEALTH & HUMAN SERVICES.  
XX  
XX Newton DL, Rybak SM;  
XX  
XX MPI: 1999-610847/52.  
XX  
XX N-PSDB; AAZ08130.  
XX  
XX New recombinant ribonucleases, used for killing target cells, e.g. for  
XX treating cancers, viral infections or autoimmune diseases  
XX

PS Claim 22; Page 62; 71pp; English.  
XX  
CC The present sequence is a Rana catesbeiana oocyte ribonuclease (RacOR1)  
CC protein encoded by a cDNA modified for expression in E. coli. Carboxy  
CC terminal end of RacOR1 has a covalently bound ligand binding moiety,  
CC which can be a LL2 antibody directed against CD22 on cancerous B cells  
CC or human chorionic gonadotropin (hCG) effective against Kaposi's  
CC Sarcoma cells. Recombinant ribonucleases can be expressed in bacteria  
CC without an N-terminal methionine due to the presence of a signal peptide  
CC that is cleaved by bacteria. The soluble expression of ribonuclease  
CC allows the proteins to be fused in-frame with ligand binding moieties to  
CC form cytotoxic fusion proteins. They can be used for treatment of cancer  
CC and autoimmune diseases.  
XX  
SQ Sequence 110 AA;  
Query Match 99.3%; Score 597; DB 20; Length 110;  
Best Local Similarity 100.0%; Pred. No. 8.9e+61;  
Matches 109; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
OY 2 NMAATFOOKHIIIMPICNTIMDNNIYIVGGCKRVNFTFISSATTVAICGVINMNVLS 61  
Db 2 NMAATFOOKHIIIMPICNTIMDNNIYIVGGCKRVNFTFISSATTVAICGVINMNVLS 61  
OY 62 TTRFOLMTCRTSITPRCPYSSRTETNYICVKCENQPVHFAIGRCP 110  
Db 62 TTRFOLMTCRTSITPRCPYSSRTETNYICVKCENQPVHFAIGRCP 110  
RESULT 4  
AY28873 ID AAY28873 standard; Protein: 111 AA.  
XX  
AC AAY28873;  
XX  
DT 25-JAN-2000 (first entry)  
XX  
DE Recombinant Met(-1) RacOR1.  
XX  
KW Recombinant Met(-1) Rana catesbeiana oocyte ribonuclease; RacOR1; CD22;  
KW covalently bound; LL2 antibody; ligand binding moiety; cancerous B cell;  
KW Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide;  
KW recombinant ribonuclease; cytotoxic fusion protein; cancer; bullfrog;  
KW RNase; autoimmune disease.  
XX  
OS Rana catesbeiana.  
OS Synthetic.  
XX  
FH Key Location/Qualifiers  
FT Misc-difference 1 /note= "Met not found in wild type RacOR1"  
FT FT  
XX  
PN MO9950398-A2.  
XX  
PD 07-OCT-1999.  
XX  
PF 26-MAR-1999; 99WO-US06641.  
XX  
PR 27-MAR-1998; 98US-0079751.  
XX  
PA (USSH ) US DEPT HEALTH & HUMAN SERVICES.  
XX  
PI Newton DL, Rybak SM;  
XX  
DR WPI: 1999-610847/52.  
DR N-PSDB: AA208131.  
XX  
PT New recombinant ribonucleases, used for killing target cells, e.g. for  
PT treating cancers, viral infections or autoimmune diseases  
XX  
PS Claim 22; Page 63; 71pp; English.  
XX  
CC The present sequence is a recombinant Rana catesbeiana oocyte

CC ribonuclease (RacOR1) protein with Met at position 1. Carboxy terminal  
CC end of recombinant RacOR1 has a covalently bound ligand binding moiety,  
CC which can be a LL2 antibody directed against CD22 on cancerous B cells or  
CC human chorionic gonadotropin (hCG) effective against Kaposi's sarcoma  
CC cells. Recombinant ribonucleases can be expressed in bacteria without an  
CC N-terminal methionine due to the presence of a signal peptide that is  
CC cleaved by bacteria. The soluble expression of ribonuclease allows the  
CC proteins to be fused in-frame with ligand binding moieties to form  
CC cytotoxic fusion proteins. They can be used for treatment of cancer and  
CC autoimmune diseases.  
XX  
SQ Sequence 111 AA;  
Query Match 99.3%; Score 597; DB 20; Length 111;  
Best Local Similarity 100.0%; Pred. No. 9e+61;  
Matches 109; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
OY 2 NMAATFOOKHIIIMPICNTIMDNNIYIVGGCKRVNFTFISSATTVAICGVINMNVLS 61  
Db 3 NMAATFOOKHIIIMPICNTIMDNNIYIVGGCKRVNFTFISSATTVAICGVINMNVLS 62  
OY 62 TTRFOLMTCRTSITPRCPYSSRTETNYICVKCENQPVHFAIGRCP 110  
Db 63 TTRFOLMTCRTSITPRCPYSSRTETNYICVKCENQPVHFAIGRCP 111  
RESULT 5  
AY28874 ID AAY28874 standard; Protein: 110 AA.  
XX  
AC AAY28874;  
XX  
DT 25-JAN-2000 (first entry)  
XX  
DE Recombinant RacOR1 Met22Leu Met57Leu amino acid sequence.  
XX  
KW Recombinant Rana catesbeiana oocyte ribonuclease; covalently bound;  
KW RacOR1 Met22Leu Met57Leu; LL2 antibody; ligand binding moiety; CD22;  
KW cancerous B cell; Kaposi's sarcoma; human chorionic gonadotropin; hCG;  
KW signal peptide; recombinant ribonuclease; cytotoxic fusion protein;  
KW cancer; bullfrog; RNase; autoimmune disease.  
XX  
OS Rana catesbeiana.  
OS Synthetic.  
XX  
FH Key Location/Qualifiers  
FT Misc-difference 22 /note= "wild type Met replaced with Leu"  
FT FT  
FT Misc-difference 57 /note= "wild type Met replaced with Leu"  
FT FT  
XX  
PN MO9950398-A2.  
XX  
PD 07-OCT-1999.  
XX  
PF 26-MAR-1999; 99WO-US06641.  
XX  
PR 27-MAR-1998; 98US-0079751.  
XX  
PA (USSH ) US DEPT HEALTH & HUMAN SERVICES.  
XX  
PI Newton DL, Rybak SM;  
XX  
DR WPI: 1999-610847/52.  
DR N-PSDB: AA208132.  
XX  
PT New recombinant ribonucleases, used for killing target cells, e.g. for  
PT treating cancers, viral infections or autoimmune diseases  
XX  
PS Claim 22; Page 64; 71pp; English.  
XX  
CC The present sequence is a recombinant Rana catesbeiana oocyte  
CC ribonuclease (RacOR1) protein with Met22Leu Met57Leu. Carboxy terminal

CC end of recombinant RacOR1 has a covalently bound ligand binding moiety,  
 CC which can be a LL2 antibody directed against CD22 on cancerous B cells  
 CC or human chorionic gonadotropin (hCG) effective against Kaposi's sarcoma  
 CC cells. Recombinant ribonucleases can be expressed in bacteria without an  
 CC N-terminal methionine due to the presence of a signal peptide that is  
 CC cleaved by bacteria. The soluble expression of ribonuclease allows the  
 CC proteins to be fused in-frame with ligand binding moieties to form  
 CC cytotoxic fusion proteins. They can be used for treatment of cancer and  
 CC autoimmune diseases.

XX Sequence 110 AA:

Query Match 98.3%; Score 591; DB 20; Length 110;  
 Best Local Similarity 98.2%; Pred. No. 4.3e-60;  
 Matches 107; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

QY 2 NMATFOQKHINPIICNTIMDNNIYVGGCKRVNTFIISATYKAICTGVINNVLS 61  
 DB 2 NMATFOQKHINPIICNTIMDNNIYVGGCKRVNTFIISATYKAICTGVINNVLS 61  
 QY 62 TTRFOLNTCTRTSITPRPCPYSSRTETNYICVCKENQYVHFAGIGRCP 110  
 DB 62 TTRFOLNTCTRTSITPRPCPYSSRTETNYICVCKENQYVHFAGIGRCP 110

RESULT 6  
 AAY28876  
 ID AAY28876 standard; Protein: 111 AA.

XX AAY28876:  
 DT 25-JAN-2000 (first entry)  
 XX  
 DE Recombinant Met(-1) RacOR1 Met22Leu Met57Leu-(His)6 protein.  
 XX  
 KW Met(-1) Rana catesbeiana ribonuclease Met22Leu Met57Leu-(His)6; RacOR1;  
 KW recombinant; CD22; covalently bound; LL2 antibody; ligand binding moiety;  
 KW cancerous B cell; Kaposi's sarcoma; human chorionic gonadotropin; hCG;  
 KW signal peptide; recombinant ribonuclease; cytotoxic fusion protein;  
 KW cancer; bullfrog; RNase; autoimmune disease.  
 XX  
 OS Rana catesbeiana.  
 OS Synthetic.

XX Key Location/Qualifiers  
 FT MISC-difference 1 /note= "(His)6 histidine tag attached to N-terminal Met"  
 FT MISC-difference 1 /note= "Met not found in wild type RacOR1"  
 FT MISC-difference 23 /note= "Wild type Met replaced with Leu"  
 FT MISC-difference 58 /note= "Wild type Met replaced with Leu"  
 FT  
 XX WO9950398-A2.  
 XX 07-OCT-1999.  
 XX PD  
 XX 26-MAR-1999; 99WO-US06641.  
 XX PF  
 XX 27-MAR-1998; 98US-0079751.  
 XX PR  
 XX (USSH ) US DEPT HEALTH & HUMAN SERVICES.  
 XX PA  
 XX Newton DL, Rybak SM;  
 XX PI  
 XX WPI: 1999-610847/52.  
 XX DR N-PSDB; AAZ08133.  
 XX DX  
 XX New recombinant ribonucleases, used for killing target cells, e.g. for  
 XX PT treating cancers, viral infections or autoimmune diseases  
 XX PS Claim 22; Page 66; 71pp; English.

XX The present sequence is a recombinant Rana catesbeiana oocyte  
 CC ribonuclease (RacOR1) protein with Met at position 1 attached to a  
 CC (His)6 tag, Met23Leu and Met58Leu. Carboxy terminal end of recombinant  
 CC RacOR1 has a covalently bound ligand binding moiety, which can be a LL2  
 CC antibody directed against CD22 on cancerous B cells or human chorionic  
 CC gonadotropin (hCG) effective against Kaposi's sarcoma cells. Recombinant  
 CC ribonucleases can be expressed in bacteria without an N-terminal  
 CC methionine due to the presence of a signal peptide that is cleaved by  
 CC bacteria. The soluble expression of ribonuclease allows the proteins to  
 CC be fused in-frame with ligand binding moieties to form cytotoxic fusion  
 CC proteins. They can be used for treatment of cancer and autoimmune  
 CC diseases.

XX Sequence 111 AA:

Query Match 98.3%; Score 591; DB 20; Length 111;  
 Best Local Similarity 98.2%; Pred. No. 4.4e-60;  
 Matches 107; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

QY 2 NMATFOQKHINPIICNTIMDNNIYVGGCKRVNTFIISATYKAICTGVINNVLS 61  
 DB 3 NMATFOQKHINPIICNTIMDNNIYVGGCKRVNTFIISATYKAICTGVINNVLS 62  
 QY 62 TTRFOLNTCTRTSITPRPCPYSSRTETNYICVCKENQYVHFAGIGRCP 110  
 DB 63 TTRFOLNTCTRTSITPRPCPYSSRTETNYICVCKENQYVHFAGIGRCP 111

RESULT 7  
 AAY33321  
 ID AAY33321 standard; Protein: 111 AA.

XX AAY33321:  
 DT 29-NOV-1999 (first entry)  
 XX  
 DE Frog lectin protein fragment.  
 XX  
 KW Cytotoxic; RNase; ribonuclease; pancreatic; antibody; light chain;  
 KW heavy chain; cell surface marker; treatment; tumor; viral infection;  
 KW parasite infection; immune dysfunctional cell; autoimmune disease;  
 KW contraceptive; cell separation; transplantation; bone marrow ablation;  
 KW leukemia cell; T-cell; graft-versus-host disease; bullfrog; lectin.  
 XX  
 OS Rana catesbeiana.  
 OS  
 XX US955073-A.  
 XX PN  
 XX 21-SEP-1999.  
 XX PD  
 XX 09-JUL-1997; 97US-0891848.  
 XX PF  
 XX 22-SEP-1993; 93US-0125462.  
 XX PR 22-OCT-1991; 91US-0779195.  
 XX PR 20-APR-1990; 90US-0510696.  
 XX PR 04-FEB-1993; 93US-0014082.  
 XX PA  
 XX (USSH ) US DEPT HEALTH & HUMAN SERVICES.  
 XX XX  
 XX Rybak SM, Newton DL, Nicholls PJ, Youle RJ;  
 XX PI  
 XX WPI: 1999-560488/47.  
 XX DR  
 XX Recombinantly fused pancreatic RNase-targeting proteins useful for  
 XX PT treating tumors, infections, immune or autoimmune disorders and as a  
 XX PT contraceptive  
 XX PS Example 3; Fig 19; 47pp; English.  
 XX PS This invention describes a novel nucleic acid construct comprising  
 CC sequences encoding functional pancreatic RNase and a second protein  
 CC (preferably the light and heavy chains of an antibody) which binds a

CC specific cell surface marker on a target cell and functions as a  
 CC cytotoxic agent. The products can be used for selectively killing cells  
 CC expressing a specific surface marker. They can be used for treating  
 CC tumors or infected cells (e.g. cells infected by viruses (especially  
 CC latent or chronic virus infections, such as human immunodeficiency virus  
 CC (HIV)-1, Epstein-Barr virus, herpes viruses (herpes simplex types 1 and  
 CC 11), hepatitis viruses (B, non-A-non-B, and delta), herpes zoster,  
 CC cytomegalovirus) and cells infected with parasites (such as the malaria  
 CC parasite)). They can also be used for treating immune dysfunctional cells  
 CC in immune and autoimmune diseases. Additionally, they may be used as  
 CC contraceptives. Finally they can also be used for cell separation in  
 CC vitro by selectively killing unwanted types of cells (e.g. in bone  
 CC marrow) prior to transplantation into a patient undergoing marrow  
 CC ablation by radiation or for killing leukemia cells or T-cells that would  
 CC cause graft-versus-host disease. This sequence represents a fullfrog  
 CC (Rana catesbeiana) lectin used to describe the method of the invention.

SQ Sequence 111 AA:

Query Match 97.6%; Score 586.5; DB 20; Length 111;  
 Best Local Similarity 99.1%; Pred. No. 1.4e-59;  
 Matches 109; Conservative 0; Mismatches 0; Indels 1; Gaps 1;

OY 2 NNATFOOKHIIINPTII-CNTIMDNNTIYIGGCKRVNFTIISATTVKATCGVINMNV 60  
 DB 2 NNATFOOKHIIINPTIINCMNTIMDNNTIYIGGCKRVNFTIISATTVKATCGVINMNV 61  
 OY 61 STTRFQNLNCTRTSTTPRCPYSSRTETNYICVKCENQPVHFGIGRC 110  
 DB 62 STTRFQNLNCTRTSTTPRCPYSSRTETNYICVKCENQPVHFGIGRC 111

RESULT 8

AAW06544  
 ID AAW06544 standard; protein; 104 AA.

AAW06544;

22-AUG-1997 (first entry)

Antitumour protein from Rana pipiens oocytes.

Tumour; chemotherapy; radiotherapy; frog.

Rana pipiens.

MO9639428-A1.

12-DEC-1996.

03-JUN-1996; 96WO-US08304.

06-JUN-1995; 95US-0467955.

(ALFA-) ALFACELL CORP.

Ardelet WJ;

WPI; 1997-043063/04

Antitumour proteins from Rana pipiens oocyte(s) - have fewer  
 disadvantages than chemotherapy, surgery and radiotherapy

Claim 8; Page 28; 45pp; English.

The present sequence is a specifically claimed example of an  
 antitumour protein from the generic protein in AAW18224, with the  
 molecular weight 12000. This is one of two preferred proteins (the  
 other in AAW06543) that have been isolated from Rana pipiens oocytes.  
 Both proteins have a blocked amino terminal group and are essentially  
 free of carbohydrates. The proteins are used to treat tumours, use of  
 the peptides has fewer disadvantages than chemotherapy, radiotherapy  
 and surgery in the treatment of tumours.

XX Sequence 104 AA:

SQ Query Match 47.0%; Score 282.5; DB 18; Length 104;  
 Best Local Similarity 50.0%; Pred. No. 1.1e-24;  
 Matches 55; Conservative 15; Mismatches 31; Indels 9; Gaps 4;

OY 2 NNATFOOKHIIIN-PIICNTIMDNNTIYIGGCKRVNFTIISATTVKATCGVI-NMNV 59  
 DB 2 DMTEFOKKHVTNRDVCNNINMSTNF-----HKDKNTFIYSPEPVKATCGIISKNV 57  
 OY 60 LSTTRFQNLNCTRTSTTPRCPYSSRTETNYICVKCENQPVHFGIGRC 109  
 DB 58 LTTSEFYLSDC---NVTSPKCKYKLTSTNKFCVTCENQAPVHFGVGR 104

RESULT 9

AAV28870  
 ID AAV28870 standard; protein; 104 AA.

AAV28870;

25-JAN-2000 (first entry)

Recombinant RapL1 GlnIser amino acid sequence.

Recombinant Rana pipiens ribonuclease; RapL1 GlnIser; covalently bound;

KL2 antibody; ligand binding moiety; CD22; cancerous B cell; frog;

Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide;

recombinant ribonuclease; cytotoxic fusion protein; cancer; Rana;

autoimmune disease.

Rana pipiens.

Synthetic.

MO9950398-A2.

07-OCT-1999.

26-MAR-1999; 99WO-US06641.

27-MAR-1998; 98US-0079751.

(USSH ) US DEPT HEALTH & HUMAN SERVICES.

Newton DL, Rybak SM;

WPI; 1999-610847/52.

N-PSDB; AA208128.

New recombinant ribonucleases, used for killing target cells, e.g. for

treating cancers, viral infections or autoimmune diseases

Claim 34; Page 60; 71pp; English.

The present sequence is a recombinant Rana pipiens ribonuclease (RapL1)  
 protein with GlnIser. Carboxy terminal end of recombinant RapL1 has a  
 covalently bound ligand binding moiety, which can be a KL2 antibody  
 directed against CD22 on cancerous B cells or human chorionic  
 gonadotropin (hCG) effective against Kaposi's sarcoma cells. Recombinant  
 ribonucleases can be expressed in bacteria without an N-terminal  
 methionine due to the presence of a signal peptide that is cleaved by  
 bacteria. The soluble expression of ribonuclease allows the proteins to  
 be fused in-frame with ligand binding moieties to form cytotoxic fusion  
 proteins. They can be used for treatment of cancer and autoimmune  
 diseases.

SQ Sequence 104 AA:

Query Match 46.7% Score 280.5; DB 20; Length 104;  
Best Local Similarity 49.5%; Pred. No. 1.8e-24;  
Matches 55; Conservative 15; Mismatches 32; Indels 9; Gaps 4;

OY 1 SMMATFOOKHILIN-PIICNTIMDNNIYIYGCGCKRVNTEFISSATYVKAICGVI-NMN 58  
Db 1 SDMLTFCKKHILNTRDVCNNIMSTNLF---HCKDKNTFIYSRPEPVKAICKGIASKN 56

OY 59 VLSSTRPOLNCTRTSITPRCPYSSRTEINVCCKENQYPVHAFAGIGRC 109  
Db 57 VLTISEEYLSDC---NNTSRCKYKLLKSTNTFCVTCENQAPVHAFVGVGHC 104

## RESULT 10

AAV28871  
ID AAV28871 standard; Protein; 105 AA.

XX AAV28871;

XX 25-JAN-2000 (first entry)

XX Recombinant Met(-1) RapLRI GlnSer amino acid sequence.

XX Recombinant Met(-1) Rana pipiens ribonuclease GlnSer; RapLRI; CD22;  
KM covalently bound; LL2 antibody; ligand binding moiety; cancerous B cell;  
KM Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide;  
KM recombinant ribonuclease; cytotoxic fusion protein; cancer; frog;  
XX autoimmune disease; RNase.

OS Rana pipiens.  
XX Synthetic.

XX Key Location/Qualifiers

FT MISC-difference 1 /note= "Met not found in wild type RapLRI"

FT MISC-difference 2 /note= "Wild type Gln replaced with Ser"

FT W09950398-A2.

PN 07-OCT-1999.

PD 26-MAR-1999; 99WO-US06641.

XX 27-MAR-1998; 98US-0079751.

XX (USSH ) US DEPT HEALTH & HUMAN SERVICES.

XX Newton DL, Rybak SM;

XX WPI; 1999-610847/52.

XX N-PSDB; AA208129.

XX New recombinant ribonucleases, used for killing target cells, e.g. for  
treating cancers, viral infections or autoimmune diseases

XX Claim 34; Page 61; 71pp; English.

XX The present sequence is a recombinant Rana pipiens ribonuclease (RapLRI)  
CC protein with Met at position 1 and GlnSer carboxy terminal end of  
CC recombinant RapLRI has a covalently bound ligand binding moiety, which  
CC can be a LL2 antibody directed against CD22 on cancerous B cells or human  
CC chorionic gonadotropin (hCG) effective against Kaposi's sarcoma cells.  
CC Recombinant ribonucleases can be expressed in bacteria without an N-  
CC terminal methionine due to the presence of a signal peptide that is  
CC cleaved by bacteria. The soluble expression of ribonuclease allows the  
CC proteins to be fused in-frame with ligand binding moieties to form  
CC cytotoxic fusion proteins. They can be used for treatment of cancer and  
CC autoimmune diseases.

XX Sequence 105 AA;

XX Query Match 46.7% Score 280.5; DB 20; Length 105;

Best Local Similarity 49.5%; Pred. No. 1.8e-24;  
Matches 55; Conservative 15; Mismatches 32; Indels 9; Gaps 4;

OY 1 SMMATFOOKHILIN-PIICNTIMDNNIYIYGCGCKRVNTEFISSATYVKAICGVI-NMN 58  
Db 2 SDMLTFCKKHILNTRDVCNNIMSTNLF---HCKDKNTFIYSRPEPVKAICKGIASKN 57

OY 59 VLSSTRPOLNCTRTSITPRCPYSSRTEINVCCKENQYPVHAFAGIGRC 109  
Db 58 VLTISEEYLSDC---NNTSRCKYKLLKSTNTFCVTCENQAPVHAFVGVGHC 105

## RESULT 11

AAV28865  
ID AAV28865 standard; Protein; 104 AA.

XX AAV28865;

XX 25-JAN-2000 (first entry)

XX Rana pipiens liver ribonuclease (RapLRI).

XX Rana pipiens liver ribonuclease; RapLRI; covalently bound; LL2 antibody;  
KM ligand binding moiety; CD22; cancerous B cell; Kaposi's sarcoma; frog;  
KM human chorionic gonadotropin; hCG; recombinant ribonuclease; RNase;  
KM signal peptide; cytotoxic fusion protein; cancer; autoimmune disease.

XX Rana pipiens.

OS W09950398-A2.

PN 07-OCT-1999.

PD 26-MAR-1999; 99WO-US06641.

XX 27-MAR-1998; 98US-0079751.

XX (USSH ) US DEPT HEALTH & HUMAN SERVICES.

XX Newton DL, Rybak SM;

XX WPI; 1999-610847/52.

XX N-PSDB; AA208124.

XX New recombinant ribonucleases, used for killing target cells, e.g. for  
treating cancers, viral infections or autoimmune diseases

XX Claim 1; Page 55; 71pp; English.

XX The present sequence is Rana pipiens liver ribonuclease (RapLRI)  
CC protein. Carboxy terminal end of RapLRI has a covalently bound  
CC ligand binding moiety, which can be a LL2 antibody directed against  
CC CD22 on cancerous B cells or human chorionic gonadotropin (hCG)  
CC effective against Kaposi's sarcoma cells. Recombinant ribonucleases can  
CC be expressed in bacteria without an N-terminal methionine due to the  
CC presence of a signal peptide that is cleaved by bacteria. The soluble  
CC expression of ribonuclease allows the proteins to be fused in-frame with  
CC ligand binding moieties to form cytotoxic fusion proteins. They can be  
CC used for treatment of cancer and autoimmune diseases.

XX Sequence 104 AA;

XX Query Match 46.0% Score 276.5; DB 20; Length 104;

XX Best Local Similarity 49.1%; Pred. No. 5.2e-24;  
Matches 54; Conservative 15; Mismatches 32; Indels 9; Gaps 4;

OY 2 SMMATFOOKHILIN-PIICNTIMDNNIYIYGCGCKRVNTEFISSATYVKAICGVI-NMN 59  
Db 2 SDMLTFCKKHILNTRDVCNNIMSTNLF---HCKDKNTFIYSRPEPVKAICKGIASKN 57

OY 60 LSTTRPOLNCTRTSITPRCPYSSRTEINVCCKENQYPVHAFAGIGRC 109  
Db 58 LVTISEEYLSDC---NNTSRCKYKLLKSTNTFCVTCENQAPVHAFVGVGHC 104



